

*Judith Keating***PHYSICS OF REACTOR SAFETY****Quarterly Report****April—June 1976**

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PHYSICS OF REACTOR SAFETY

Quarterly Report  
April—June 1976

Applied Physics Division

Work performed for the  
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Previous reports in this series

ANL-75-67	April—June 1975
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ANL-76-13	October—December 1975
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## PHYSICS OF REACTOR SAFETY

Quarterly Report  
April-June 1976

## I. ABSTRACT

This quarterly progress report summarizes work done in Argonne National Laboratory's Applied Physics Division for the Division of Reactor Safety Research of the U.S. Nuclear Regulatory Commission during the months of April-June 1976. It includes reports on reactor safety research and technical coordination of the RSR safety analysis program by members of the Reactor Safety Appraisals Group, Monte Carlo analysis of safety-related critical assembly experiments by members of the Theoretical Fast Reactor Physics Group, and planning of DEMO safety-related critical experiments by members of the Zero Power Reactor (ZPR) Planning and Experiments Group.

TECHNICAL COORDINATION - FAST REACTOR  
SAFETY ANALYSIS  
(A2015)

## II. SUMMARY

Comparisons of reactivity coefficients and power distributions for the BOL state of the CRBR revealed that the results obtained assuming LWR-grade plutonium generally differed little from those obtained assuming FFTF-grade plutonium. A major part of the difference in reactivity worths measured in dollars arises from a 9% difference in  $\beta_{\text{eff}}$  calculated for the two cases.

Comparison of reactivity coefficients for the EOEC state of the CRBR using ENDF/B Version III and Version IV indicated generally minor differences, except that the positive sodium void effect over the core increased by 20 to 25%.

Comparison of reactivity coefficients for the CRBR using R-Z and two-dimensional hexagonal mesh calculations indicated that the most significant errors in the R-Z model were in the sodium and steel worths. Errors in the sodium worth in a considerable part of the core were in the range of 20-40% for the BOEC case. For the EOEC case, withdrawal of control rods resulted in smaller flux variations and therefore smaller errors in the R-Z model.

It was determined that the net effect of resonance self-overlap on the CRBR Doppler effect was about 8%, so that uncertainty in treatment of this effect should be less than 8%.

A new FX2-POOL computational path has been written to calculate the potential autocatalytic effect of bubble collapse in a boiling fuel-steel pool.

POOL and FX2-POOL were modified to calculate the work energy available from fuel and steel at any time during an HCDA. The effect of fuel to steel heat transfer has been found to have a negligible effect on the total energy release in an HCDA, but can have an important effect on work energy on a time scale of 20 to 30 ms. Fuel-steel heat transfer should reduce the work energy as the steel would have to have an initial temperature of more than 2500°K for the work energy to be increased.

Programming of the multinode failure option in the EPIC fuel-coolant interaction code and of capability for a dealing with a partially voided channel has been completed. The first version of the code is now in a final testing phase to be sure that all options operate successfully.

It is hoped to have a preliminary plan for validation of the SIMMER and HAARM codes by the end of calendar year 1976. Strong areas of mutual benefit have been identified between the HAARM experimental effort and the SIMMER analysis.

### III. STUDY OF BASIC PROBLEMS IN ACCIDENT ANALYSIS

#### A. Initiating Condition Variations

##### 1. Power and Material Worth Distribution in the FFTF-Grade Plutonium-Fueled Clinch River Breeder Reactor at the Beginning of Life (Kalimullah and H. H. Hummel)

We have summarized here the results of some calculations of power, sodium void, steel and core fuel worth distributions for the FFTF-grade plutonium fueled CRBR at BOL and have compared these results with those for the LWR-grade plutonium fueled CRBR at BOL reported earlier.<sup>1</sup> The isotopic composition of the FFTF-grade plutonium,  $^{238}\text{Pu}/^{239}\text{Pu}/^{240}\text{Pu}/^{241}\text{Pu}/^{242}\text{Pu}$ , is 0.0/86.4/11.7/1.7/0.2 w/o compared to 1.0/67.3/19.2/10.1/2.4 w/o of the LWR discharge grade plutonium. With the FFTF-grade plutonium the required enrichments of the inner and outer core regions are 17.7 and 25.6 w/o Pu/(U+Pu) compared to 18.7 and 27.1 w/o Pu/(U+Pu) for the LWR-grade plutonium fuel. The full-height hot full power R-Z model used is based on (U,Pu)<sub>2</sub>O<sub>3</sub> masses reported in the PSAR<sup>2</sup> (Table 4.3-32) and dimensions and volume fractions reported in Table D4-1, with the sodium density varying axially as a function of its steady state temperature calculated by the SAS Code.<sup>3</sup> Control rod insertions, cross-sections and other details of the calculations are the same as those used in the case of the LWR-grade plutonium. Table I summarizes the regional totals of power, sodium void, steel and core fuel worths both for the FFTF-grade and the LWR-grade plutonium fuels, the method of analysis being alike in each case. Figure 1 shows the subassembly power factors obtained from a 2-D triangular mesh diffusion theory calculation with the control rod and the 6 rods at flats of row 7 fully inserted. Figure 2 shows the axial distribution of power in all the rows of the core obtained from a 2-D R-Z diffusion theory calculation with partially inserted control rods, and with the fuel assumed at a uniform temperature of 1100°K. Comparison of regional power totals, subassembly power factors, subassembly peak-to-average power density ratios and axial power profiles between the FFTF-grade and the LWR-grade plutonium fuels shows that the power distribution remains essentially unchanged.

TABLE I. Comparison of Power and Material Worths by Region Between LWR-Pu-Fueled and FFTF-Pu-Fueled CRBR at the Beginning of Life

Power or Worth	Fuel Type	Inner Core	Outer Core	Lower Blanket	Upper Blanket	Radial Blanket	Total
Power, MWt	FFTF	531.5	404.5	7.7	6.3	25.0	975.0
	LWR	531.2	404.8	7.7	6.3	25.0	975.0
Sodium void worth, $10^3 \Delta k/k$	FFTF	11.83	-2.05	-1.03	-0.638	-1.49	6.63
	LWR	10.93	-2.37	-1.06	-0.656	-1.51	5.34
Worth of half steel, $10^3 \Delta k/k$	FFTF	-18.61	-1.15	1.18	0.829	2.08	-15.68
	LWR	-18.29	-1.05	1.18	0.832	2.09	-15.23
Core fuel worth, $10^3 \Delta k/k$	FFTF	263.8	187.5	39.58	29.64	5.92 <sup>a</sup>	526.5
	LWR	262.8	187.8	39.91	29.83	5.94 <sup>a</sup>	526.3

<sup>a</sup>Radial blanket fuel in radial blanket.

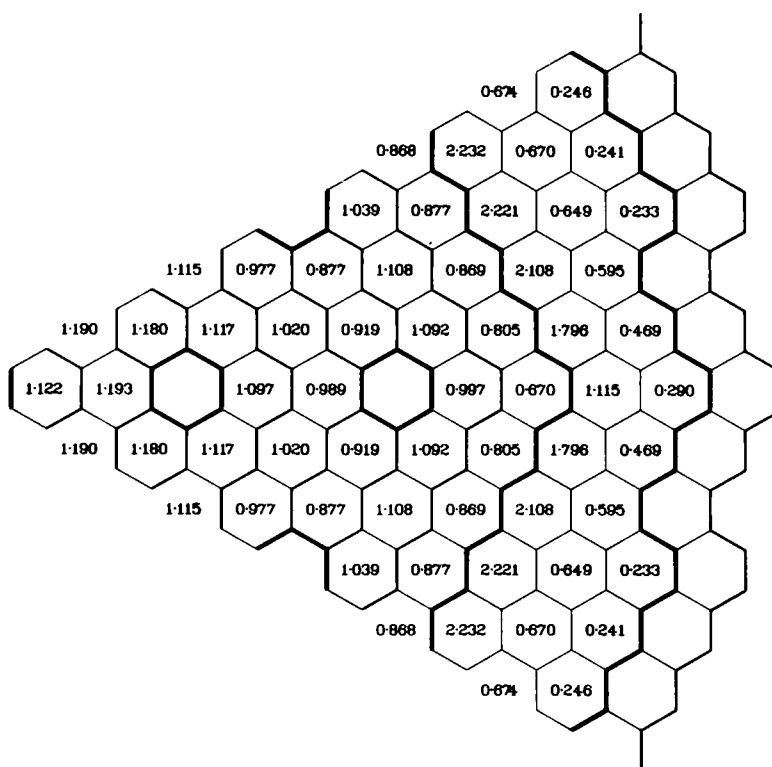


Fig. 1. Subassembly Power Factors for the FFTF Plutonium Fueled Clinch River Breeder Reactor at the Beginning of Life. ANL Neg. No. 116-76-255.

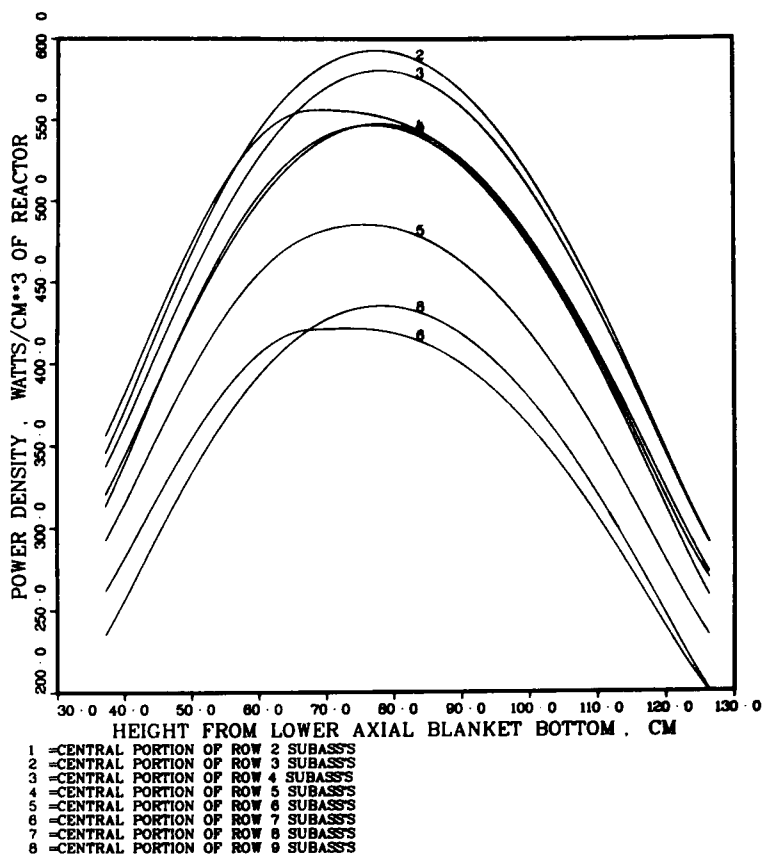


Fig. 2. Axial Distribution of Power in the FFTF-Plutonium Fueled Clinch River Breeder Reactor Core at the Beginning of Life. ANL Neg. No. 116-76-252.

The sodium void and steel worth distributions have been computed by 2-D R-Z diffusion theory linearized leakage perturbations using the real and adjoint fluxes generated in the normal reactor at a uniform fuel temperature of 1100°K. Figures 3 and 4 show comparisons of typical axial distributions in core of sodium void and steel worths for the FFTF-grade and the LWR-grade plutonium fuels (obtained by precisely the same method of analysis). In the blanket regions the comparisons are even closer than those in the core regions both for the sodium void and the steel worth distributions. When converted into dollars, (total effective delayed neutron fraction is 0.00335 using FFTF fuel compared to 0.00365 using LWR fuel as reported in the PSAR,<sup>2</sup> Table 4.3-33) the inner core sodium void worth with the FFTF-plutonium fuel is found about 18% larger than that with the LWR-plutonium fuel, whereas the corresponding comparison reported in the PSAR,<sup>2</sup> Table 4.3-35 shows an increase of about 24%. A similar comparison of the inner core steel worth shows an increase in magnitude of a little less than 11% only. Figure 5 shows a typical axial distribution of core fuel worth in all the rows of the core computed by 2-D R-Z diffusion theory linearized leakage perturbation using the real and adjoint fluxes generated in the core-and-blanket-voided reactor at a uniform fuel temperature of 1100°K. Comparison of fuel worth distributions between the FFTF-plutonium and the LWR-plutonium fueled CRBR shows that these are

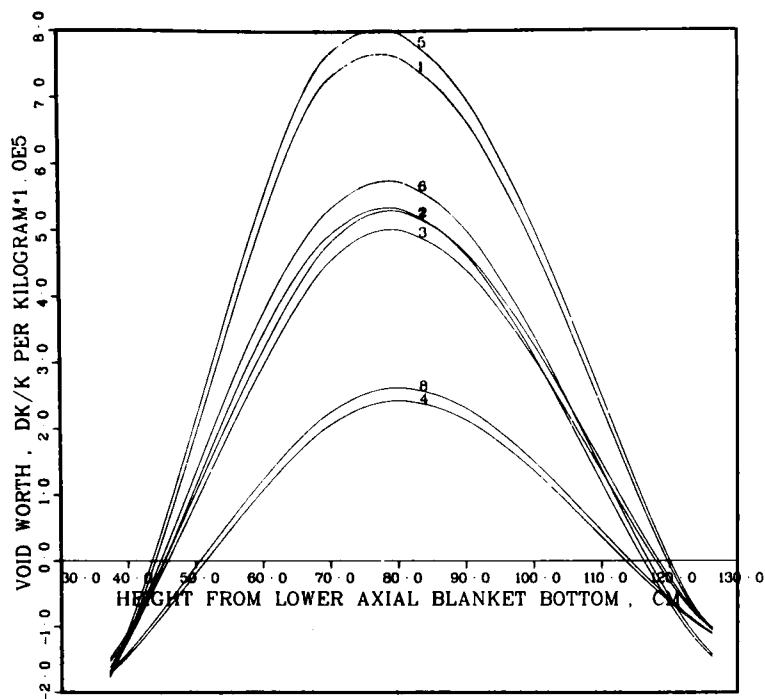


Fig. 3

Comparison of Void Worth Distributions in Core Between LWR-Pu-Fueled and FFTF-Pu-Fueled CRBR at the Beginning of Life. ANL Neg. No. 116-76-253.

- 1 = LWR-PU FUEL . CENTRAL PORTION OF ROW 2 SUBASS'S
- 2 = LWR-PU FUEL . CENTRAL PORTION OF ROW 4 SUBASS'S
- 3 = LWR-PU FUEL . CENTRAL PORTION OF ROW 6 SUBASS'S
- 4 = LWR-PU FUEL . CENTRAL PORTION OF ROW 8 SUBASS'S
- 5 = FFTF-PU FUEL . RADIAL POSITION SAME AS 1
- 6 = FFTF-PU FUEL . RADIAL POSITION SAME AS 2
- 7 = FFTF-PU FUEL . RADIAL POSITION SAME AS 3
- 8 = FFTF-PU FUEL . RADIAL POSITION SAME AS 4

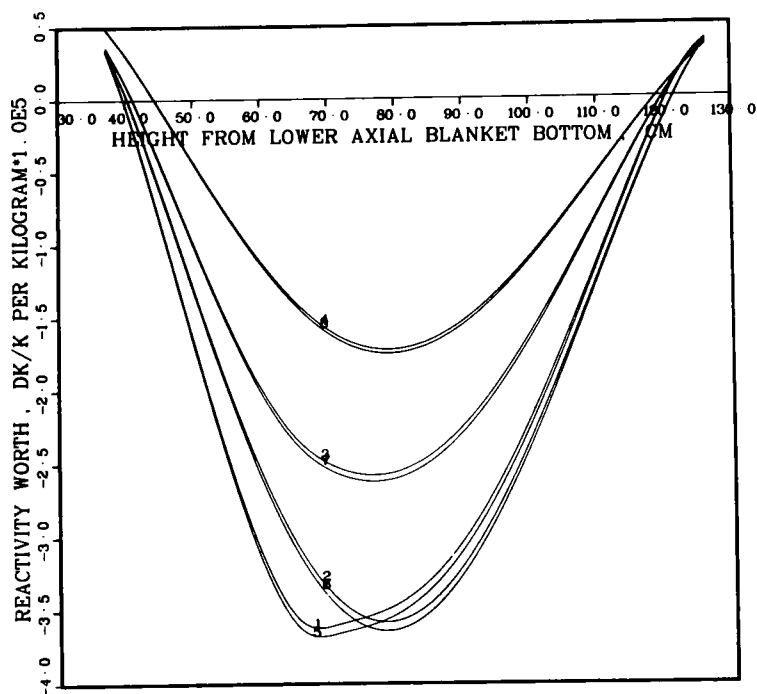


Fig. 4

Comparison of Steel Worth Distributions in Core Between LWR-Pu-Fueled and FFTF-Pu-Fueled CRBR at the Beginning of Life. ANL Neg. No. 116-76-254.

- 1 = LWR-PU FUEL . CENTRAL PORTION OF ROW 2 SUBASS'S
- 2 = LWR-PU FUEL . CENTRAL PORTION OF ROW 4 SUBASS'S
- 3 = LWR-PU FUEL . CENTRAL PORTION OF ROW 6 SUBASS'S
- 4 = LWR-PU FUEL . CENTRAL PORTION OF ROW 8 SUBASS'S
- 5 = FFTF-PU FUEL . RADIAL POSITION SAME AS 1
- 6 = FFTF-PU FUEL . RADIAL POSITION SAME AS 2
- 7 = FFTF-PU FUEL . RADIAL POSITION SAME AS 3
- 8 = FFTF-PU FUEL . RADIAL POSITION SAME AS 4

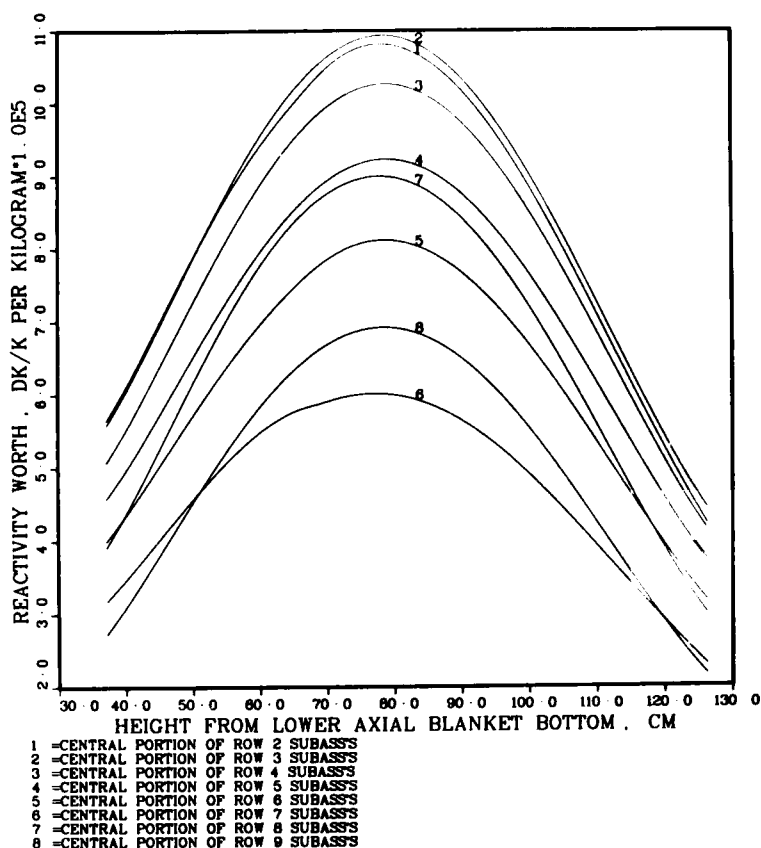


Fig. 5

Axial Distribution of Core Fuel Worth in Core in the FFTF-Plutonium Fueled Clinch River Breeder Reactor at the Beginning of Life. ANL Neg. No. 116-76-251.

essentially the same in units of  $\Delta k/k$  per kilogram. It should be pointed out that the FFTF-plutonium fuel is about 9% higher in specific reactivity worth than the LWR-plutonium fuel because of the difference in the total delayed neutron fraction.

## 2. Comparison of Reactivity Coefficients for the CRBR at End-of-Equilibrium Cycle (EOEC) as Computed with ENDF/B-III and ENDF/B-IV Based Cross Sections (P. H. Kier)

To provide information on the sensitivity of calculated reactivity coefficients to basic neutron data and procedures for processing these data into broad group cross section, reactivity coefficient computed for EOEC conditions with use of a cross section library based on ENDF/B-III data<sup>4</sup> have been recalculated with a new library based on ENDF/B-IV data. This new library has several refinements not usually made. The temperature dependence of scattering cross sections is accounted for, the constituents of stainless steel are treated as resonant materials, and integral transport theory rather than the narrow resonance approximation is used to compute resonance absorption in the resolved resonance energy range.

The results of this study will now be summarized. The use of the new library results in negligible changes in core fuel worth and power distributions. It did result, however, in significant increases in sodium void worths of about 30% (20 to 25% over the core) and increases in steel worth of about 5%. There was also an increase in the fuel Doppler coefficient of between 1 and 5%. Since the constituents of steel were treated as resonant materials in generating the new library, they have temperature-dependent cross sections and

their contribution to the Doppler effect can be calculated. With the assumption that the change in temperature in the steel is the same as in the fuel, for the compositions in CRBR at EOEC, including the Doppler effect in the constituents of steel would increase the overall Doppler coefficient by about 10%.

### 3. Comparison of Reactivity Coefficients for CRBR as Computed with the R-Z Model and with the Two-Dimensional Hexagonal Mesh Model (P. H. Kier)

It is of interest to compare reactivity coefficients as computed with two-dimensional R-Z and hexagonal-mesh perturbation theory models to gain insight into the uncertainties introduced into the calculation of these quantities from attempting to represent a three-dimensional system with two-dimensional models.

For Beginning-of-Equilibrium Cycle (BOEC) conditions for which there are partly inserted control rods, reactivity coefficients over the entire height of the core were compared for the reactor at 1100°K and with sodium unvoided. The results of this comparison are given by channel in Table II. From the table it is seen that the Doppler worth and core fuel worth are relatively insensitive to the two-dimensional representation of the system. For the steel worth, the discrepancies are larger and for the sodium void worth the discrepancies are sizable for several channels. For the sodium void worth the net reactivity as computed with first order perturbation theory was decomposed into components. Generally, the major contributions to the discrepancy arose from differences in two components: net scattering and radial leakage. There were large relative differences in the radial leakage component which indicates that local flux gradients are not well represented in two-dimensional models. Although the relative differences in the net scattering component are smaller, the magnitude of the component is larger than the radial leakage component and generally has a larger contribution to the sodium worth discrepancy.

TABLE II. Comparison of Reactivity Coefficients for Full Core Height, by Channel, for CRBR at BOEC as Obtained from 2-D Hexagonal-Mesh and R-Z Perturbation Calculations,  $\Delta k/k^2 \times 10^3$

Channel	Subass'y. I.D.	Sodium Void		Steel Removal		Doppler Worth		Core Fuel Worth	
		Hex	R-Z	Hex	R-Z	Hex	R-Z	Hex	R-Z
1	1,7	1.758	1.849	5.423	5.748	-0.3917	-0.4399	42.11	42.84
2	3,6	1.684	1.774	5.482	5.656	-0.3886	-0.4257	32.74	32.80
3	2,10,15	2.271	2.636	7.147	7.987	-0.4867	-0.5465	36.70	36.18
4	5,8,16	1.647	2.252	5.221	5.992	-0.3824	-0.3554	37.10	36.74
5	4,9,11,14	1.999	2.765	5.669	7.086	-0.4541	-0.4324	57.82	57.54
6	12,13,17	1.586	1.798	4.307	4.719	-0.2479	-0.2577	26.64	26.64
7	19,22	0.461	0.244	1.825	1.230	-0.1040	-0.1015	24.68	23.40
8	18,24,28,31,68	0.299	0.573	3.080	3.662	-0.2102	-0.2436	61.56	68.89
9	20,23,32	0.274	0.471	1.661	2.338	-0.1386	-0.1552	29.11	31.34
10	21,25,26,27,29,30	-0.643	-0.585	0.053	0.117	-0.2474	-0.2672	49.93	52.16

For End-of-Equilibrium Cycle (EOEC) conditions, for which all control rods were withdrawn from the core, reactivity coefficients were compared for a height of 5.6 cm centered about the core midplane. This comparison gives an indication of the errors introduced by azimuthal smearing of subassemblies with different compositions. For this case, the agreement between the R-Z and the hexagonal-mesh calculations were excellent for the Doppler worth and the core fuel worth and were generally significantly better for the sodium void worth and steel worth than for the BOEC case as can be seen from Table III. The significantly better agreement for the EOEC case was to be expected because the BOEC case presented a far more severe test of two-dimensional models.

TABLE III. Comparison of Reactivity Coefficients about the Midplane, by Channel, for CRBR with Normal Sodium at EOEC as Obtained from 20D Hex and R-Z Perturbation Calculations,  $\Delta k/k^2 \times 10^4$

Channel	Subass'y. I.D.	Sodium Void		Steel Removal		Doppler Worth		Core Fuel Worth	
		Hex	R-Z	Hex	R-Z	Hex	R-Z	Hex	R-Z
1	1,7	2.171	2.317	5.953	6.196	-0.3314	-0.3319	24.25	25.34
2	2,3,6	3.420	3.605	8.938	9.194	-0.4283	-0.4221	26.51	27.38
3	4,11,14	2.631	2.658	7.190	7.278	-0.4040	-0.3814	31.32	31.63
4	8,10,15,13,16	4.816	5.169	12.805	13.423	-0.6063	-0.6205	38.26	39.61
5	18,24	0.788	1.105	28.58	3.455	-0.1598	-0.1661	26.64	28.80
6	19,22	0.438	0.521	1.758	1.930	-0.1151	-0.1144	18.68	19.36
7	5,9,21,26,68	2.283	2.494	7.190	7.659	-0.4280	-0.4397	43.12	44.54
8	20,23,25	0.619	0.714	2.512	2.713	-0.1654	-0.1615	25.29	25.38
9	12,17,32	1.557	1.496	4.652	4.641	-0.2434	-0.2392	20.22	20.76
10	27,28,29,30,31	-0.414	-0.752	0.946	-0.563	-0.1802	-0.1732	34.56	35.39

#### 4. Effect of the Self-Overlap Correction on Doppler Effect Calculations (P. H. Kier)

As part of our continuing program to determine uncertainties in the computation of reactivity coefficients, a study was made of the effect of the self-overlap correction to resonance cross sections in the unresolved resonance energy range on computed Doppler reactivities. Changes in the capture cross sections of  $^{238}\text{U}$  and in the capture, fission and nu-fission cross sections of  $^{239}\text{Pu}$  between 1100 and 4400°K were obtained both including and neglecting the self-overlap correction. These two sets of Doppler difference cross sections were input as perturbations for a simplified, one-dimensional representation of the unvoided CRBR. For  $^{238}\text{U}$ , neglect of the self-overlap correction resulted in an increase in the contribution from the unresolved resonance energy range of about 25%. However, as only about a quarter of the  $^{238}\text{U}$  Doppler effect comes from the unresolved resonance energy range, the net effect was about 8%. For  $^{239}\text{Pu}$ , whose unresolved resonance energy range extends down to a few hundred electron-volts, neglect of the self-overlap correction results in an increase in the Doppler effect of about 25%; however, the contribution to the overall Doppler effect from  $^{239}\text{Pu}$  is only a few percent of  $^{238}\text{U}$  and of opposite algebraic sign. Thus it appears that any uncertainty introduced into the calculation of the Doppler effect from the treatment of the self-overlap effect would be less than 8%.



## B. Model Studies

### 1. Recriticality (P. B. Abramson)

We are continuing work on a combined compressible/incompressible flow algorithm in order to be able to follow long time (tenths of seconds) flows in pulsed pools.

We are, at this point, able to examine the influence of local pressure pulses on pools which are near  $k=1$  to study the parametric impacts on energy deposition. A new FX2-POOL calculational path will have to be written which allows POOL to call FX2 rather than vice versa as in the current FX2-POOL version. When this is accomplished, the cost of these proposed parametric studies can be reduced by as much as an order of magnitude by performing the more expensive neutronics calculations only when they are needed. We will proceed with examination of such phenomena when the new path is written.

### 2. Autocatalysis from Bubble Collapse in Pools (P. B. Abramson and T. A. Daly)

Using Monte Carlo calculations of the effect of bubble size and void fraction on diffusion coefficients performed by R. Lell and E. Gelbard, we have written a new FX2-POOL path which can calculate the autocatalytic effect of decreased neutronic streaming due to bubble collapse in a boiling pool undergoing rapid heating (and therefore liquid phase expansion) due to a prompt critical excursion. This code is now in the debugging stage, and calculations should be made in the next quarter.

### 3. Heat Transfer Effects on Work Energy (P. B. Abramson)

POOL and FX2-POOL were modified to calculate the work energy available (thermodynamic potential energy) from either fuel or steel or both, at any point along the time development of the HCDA. This was accomplished by incorporating into these codes the VENUS adiabatic expansion models used in the assessment of work energy by VENUS. Using the 1000 MW(e) Reactor calculation which we used to compare VENUS and FX2-POOL results,<sup>5</sup> calculations of the importance of fuel to steel heat transfer both during the prompt burst and in the 20 to 30 ms following the prompt burst. It was found that the fuel to steel heat transfer has essentially no role during the prompt burst (for these large ramp rate calculations of \$50/second and up). However, during the time it takes for a vapor bubble to grow to the size of the original active core (20 or 30 ms), we find that fuel to steel heat transfer can play a significant role.

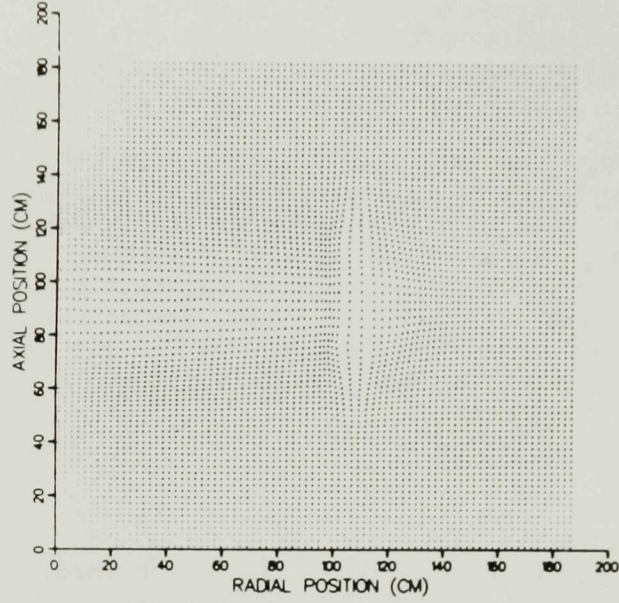
If the fuel to steel heat transfer is represented parametrically by

$$\dot{Q} = CA(T_{\text{fuel}} - T_{\text{steel}}) + A\sigma\epsilon(T_{\text{fuel}}^4 - T_{\text{steel}}^4) \quad (1)$$

then variation of C and A over a reasonable range of values can produce significant changes in the core materials behavior. Figure 6 shows the core marker particle distributions at the end of the prompt burst (6+ms) and at the time the core materials reach the blankets (30+ms).

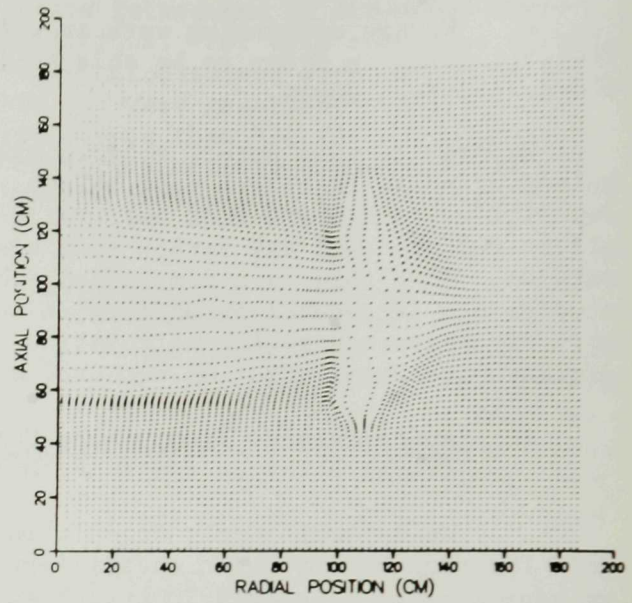
TIME(SECONDS) = 0.006360

PARTICLE DISTRIBUTION FROM HCDA



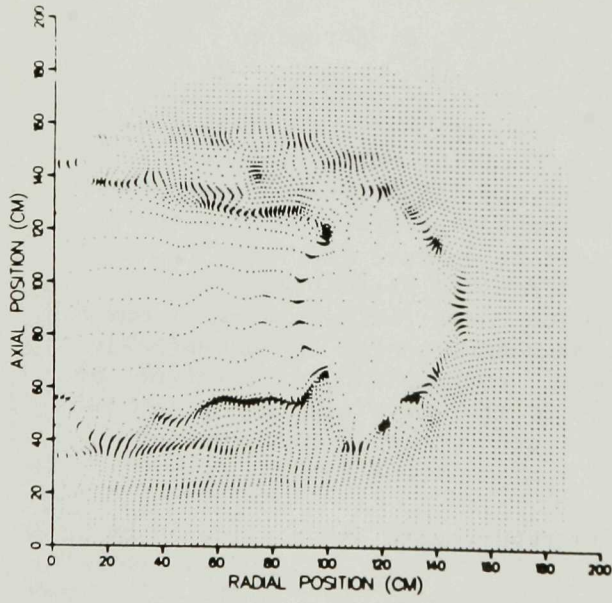
TIME(SECONDS) = 0.009860

PARTICLE DISTRIBUTION FROM HCDA



TIME(SECONDS) = 0.020360

PARTICLE DISTRIBUTION FROM HCDA



TIME(SECONDS) = 0.030360

PARTICLE DISTRIBUTION FROM HCDA

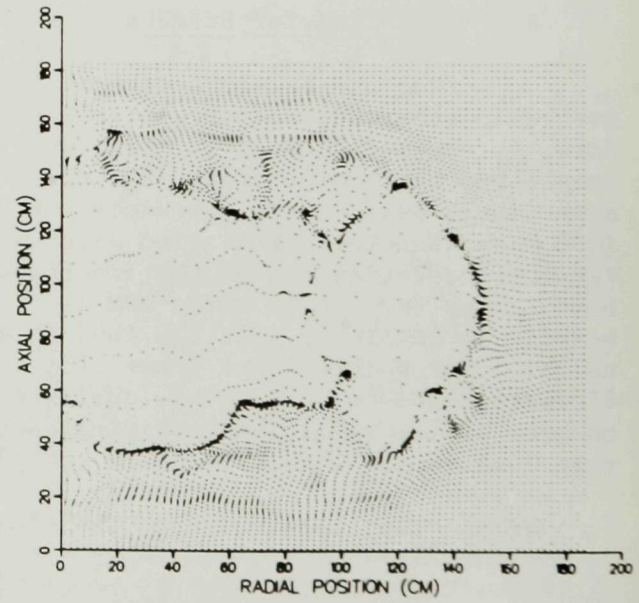


Fig. 6. Particle Configurations.  
ANL Neg. No. 116-76-125.

Figure 7 shows the influence of variation of C and A on the peak fuel temperature during that period, while Fig. 8 shows the same effects on peak pressure.

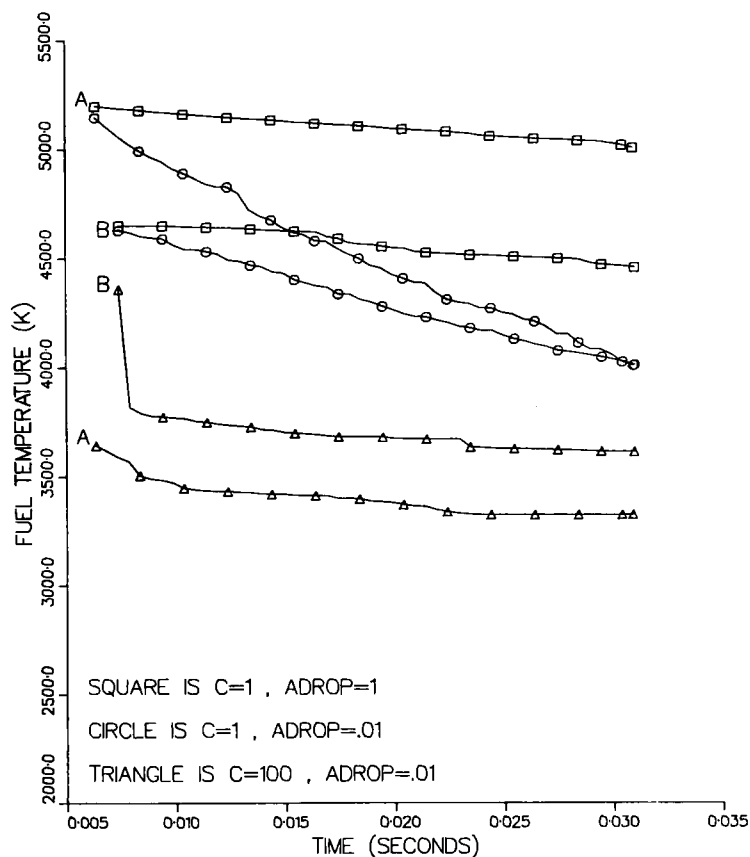


Fig. 7

Heat Transfer Effects on  
Fuel Peak Temperature.  
ANL Neg. No. 116-76-120.

Figure 9 shows how fuel to steel heat transfer can serve to reduce the available system work energy even by the time the core bubble reaches the blanket. For the calculations shown in this figure, the steel was initially at 1000°K. No work was generated from the steel in these calculations, even when all the energy was assumed to go into clad and wire wrap steel, ignoring can wall steel. Parametric variation of initial steel temperature has indicated that when the initial steel temperature is higher, additional work energy can be obtained from the steel, with the breakeven point being somewhere around 2500°K for an initial steel temperature. Furthermore, the influence of can wall steel as an energy sink serves to lower the work energy.

These studies indicate the need for further examination of the heat transfer between fuel, steel and coolant following a prompt burst. To date, damage estimates made with REXCO simply use a P-V relationship for the core bubble as a source to drive the sodium to impact. Furthermore, that P-V relationship is obtained by examination of the available work energy in fuel alone at the end of the prompt burst. Our studies indicate that such a P-V curve may be significantly in error, even when the bubble is fairly small, due to energy transfer among the core materials. The time scale for bubble growth appears to be long enough for fuel to steel and to coolant (in the LOF driven TOP) to be able to play a major role. A paper on this effect has been accepted

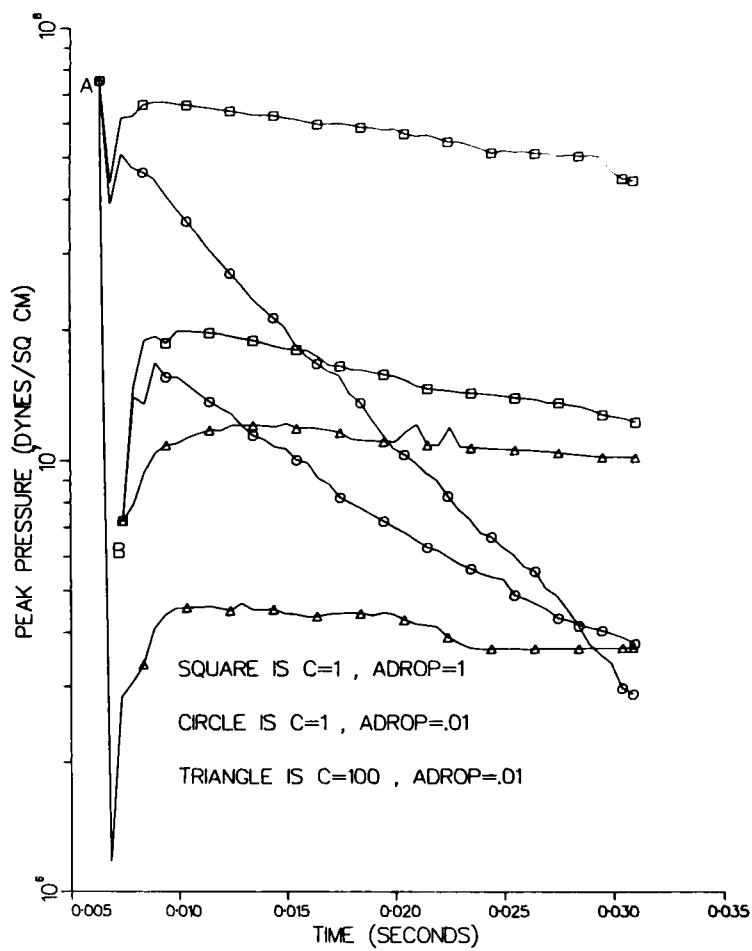


Fig. 8. Heat Transfer Effects on Peak Pressure. ANL Neg. No. 116-76-121.

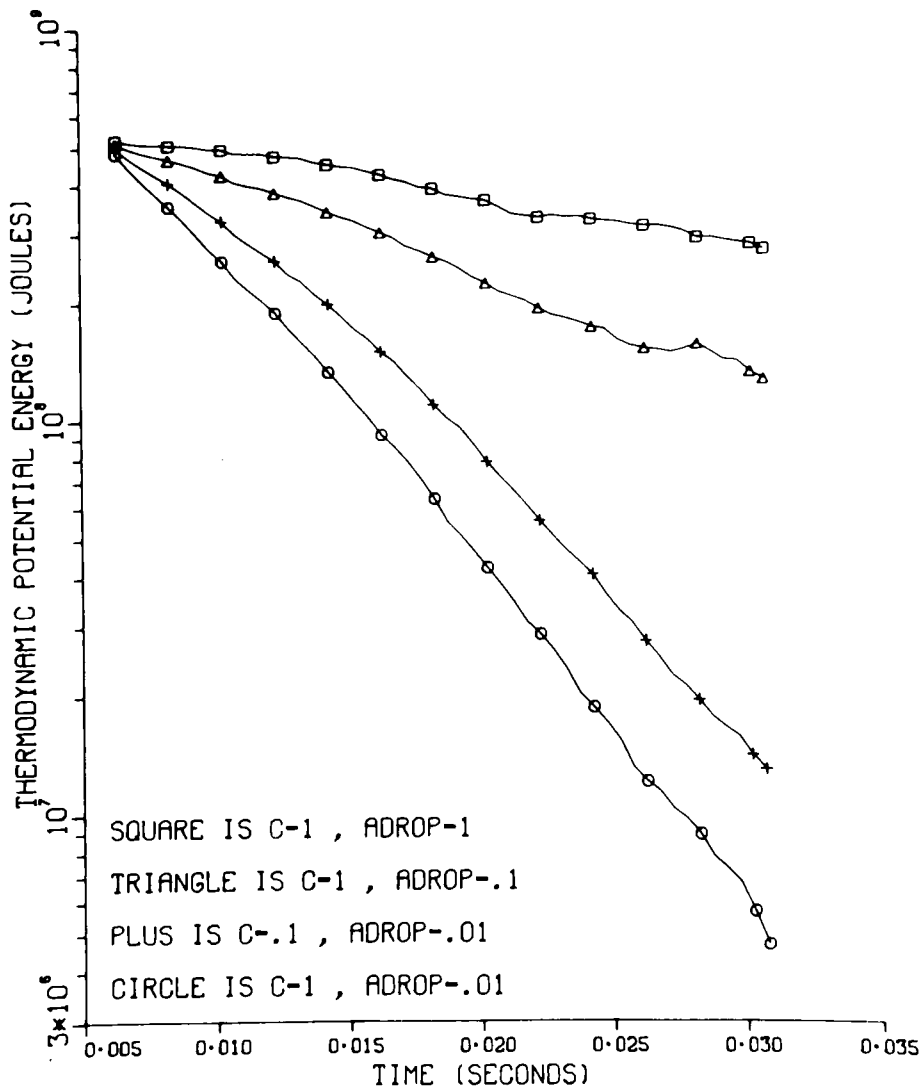


Fig. 9. Heat Transfer Effect on Work Potential.  
 ANL Neg. No. 116-76-123.

for publication at the Winter ASME meeting, and some information will also be presented at the October, 1976 Fast Reactor Safety meeting in Chicago.

#### 4. Development of EPIC (P. A. Pizzica and P. B. Abramson)

Much of the present quarter was spent refining the EPIC fuel-coolant interaction code introducing such features as a more precise tracking of the liquid sodium slug interfaces, debugging minor problems, writing an input processor, adding auxiliary editing, and breaking up the code into modules to facilitate code development and debugging. The details are unimportant, but the effort was necessary.

The multi-node failure option which will allow EPIC to treat long clad failures is working both for the case of a length fixed in time (a rip up to 25 cm long was treated) and for a rip whose length increased with time (one case increased the rip by 10 cm after a couple of msec, and then by another 10 cm after a few more msec).

The case of ejecting fuel into a partially voided coolant channel is working and stable (40 and 60% void fractions were used). Also, a multiple-node failure into the partially voided channel was successfully run. Both of these options are now undergoing checkout and comparison. When more cases have been run, a summary of the results will be presented.

Also, we intend to run a series of cases, varying the most significant input parameters such as failure location, void fraction in channel, fuel temperature, etc., in order to compare EPIC with PLUTO1. We will also use all of EPIC's features in many different cases in order to test it thoroughly before using it for production calculations.

### IV. COORDINATION OF RSR SAFETY RESEARCH

P. Abramson and H. Hummel attended an RSR meeting on SIMMER in Germantown on April 11 and a code validation meeting on April 12 in Germantown. As a result of the latter meeting, Abramson has now had two rounds of contacts with NRC ARSR consultants regarding experimental validation of SIMMER, HAARM and SSC. At this point we have received preliminary suggestions from each organization regarding their own potential contributions and are now awaiting the assessment of each reply by the other organizations. We shall attempt to have a preliminary plan on validation of SIMMER and HAARM ready by the end of calendar year 1976. Furthermore, as a result of the initial meeting, areas of strong mutual benefit have been identified between the HAARM experimental effort and the SIMMER analysis, and a cooperative effort has begun.

P. Abramson visited A.I. on April 22 to meet with R. T. Lancet and H. Morewitz to discuss recriticality and some fuel/steel interaction experiments being performed at A.I.

P. Abramson visited V. J. Dhir at UCLA on April 23 to discuss the UCLA experiments on crust formation (applied to fuel/steel pools) and to discuss debris bed experiments.

P. Abramson visited LASL on April 25 to discuss SIMMER validation analysis of transition phase, and coordination with Sandia, and visited Sandia on April 27. As a result of these meetings, several analysts from LASL visited Sandia to discuss the experimental capabilities, and a return visit is planned.

P. Abramson attended a PAHR review group on May 10 as a consultant to NRC (R. Wright), and submitted a critique of the meeting. H. Hummel, P. Abramson and P. Pizzica made presentations to the British delegation on June 24 at ANL.

## V. EVALUATION OF PROGRESS IN REACTOR SAFETY RESEARCH

Papers published in Transactions of American Nuclear Society, 23:

"A Numerical Hydrodynamics Treatment of Fuel/Steel Pools with Density Variations from Nearly Pure Vapor to Incompressible Liquid."

P. B. Abramson, p. 192.

"The Importance of Fuel to Steel Heat Transfer in Neutronic and Work Energy Release in HCDA Calculations for LMFBRs."

P. B. Abramson, p. 327.

"Power and Reactivity Distribution in the Clinch River Breeder Reactor at the Beginning-of-Life (BOL)."

Kalimullah and H. H. Hummel, p. 570.

### MONTE CARLO ANALYSIS AND CRITICALS PROGRAM PLANNING FOR SAFETY-RELATED CRITICALS (A2018)

## VI. MONTE CARLO ANALYSIS OF SAFETY-RELATED CRITICALS

As previously reported, the Monte Carlo eigenvalue for an R-Z model of the reference core agreed well with a corresponding diffusion theory eigenvalue. On the other hand, the Monte Carlo and diffusion theory eigenvalues disagreed by 1% for the step 5 damaged core. Computed eigenvalues are summarized in Table IV.

It is probably not surprising that Monte Carlo and diffusion theory eigenvalues disagree from the damaged core; it is surprising, however, that TWOTRAN disagrees so sharply with Monte Carlo and diffusion calculations for the reference core.

In order to make sure that the cross sections in TWOTRAN and DIF2D were consistent, infinite medium eigenvalues (i.e.,  $k_{\infty}$ 's) were computed for both the core and blanket materials, using both codes, with the core cross sections

TABLE IV

	Computed Eigenvalues	
	Reference Core	Damaged Core (Step 5)
Diff. Theory	1.000	1.000
VIM	$0.998 \pm 0.002^a$	$1.013 \pm 0.002^a$
TWOTRAN	1.010	Not yet available

<sup>a</sup>Confidence interval is standard deviation

as in the original R-Z calculations. It was found that TWOTRAN and DIF2D  $k_{\infty}$ 's were essentially identical. Further, after a good deal of visual checking of data sets, no cross section discrepancies have been found.

Further checking computations have been seen in spherical geometry, pressuring the R-Z core volume. Computed eigenvalues in this geometry are listed in Table V.

TABLE V

Computed Eigenvalues for 1-D Spherical Model	
Diff. Theory	1.0040
ANISN	1.0135
VIM	$1.013 \pm 0.002^a$ (100,000 histories)

<sup>a</sup>Confidence interval is standard deviation

It will be noted that in the spherical model, one finds, again that the multigroup transport (i.e., ANISN) eigenvalue is about 1% higher than the multigroup diffusion eigenvalue. This is actually, perfectly plausible in view of the very high leakage from the core, a core with  $k_{\infty} = 1.539$ . Since TWOTRAN and ANISN results show the same behavior, there is no reason to suspect any error in the TWOTRAN computation.

At this point we see two possible explanations for the VIM-TWOTRAN discrepancy in the reference core. First, it is possible that the VIM calculation, in spite of appearances, has not yet converged. Secondly, it may be that there are in the corners of the R-Z core, spectrum effects which are not adequately modelled in 27 groups. Both possibilities are being investigated. In either case the original VIM-DIF2D agreement would be fortuitous.

## VII. PLANNING OF DEMO SAFETY RELATED EXPERIMENTS

This quarterly report includes a status report on the development of the program plan for the safety related critical experiments. The initial drafts of the planned sequence of critical assembly core configurations and the related critical assembly measurements are included in Tables VI and VII.



TABLE VI. Sequence of Critical Assembly Core Configurations for  
Safety-Related Critical Experiments

Phase	Step	Approximate $k_{eff}$ of System	Measurements at Critical (See Table II)
I. Reference Core.	A. Reference Case.	1.0	Yes
	B. Measure Worth of Central B <sub>4</sub> C Control Rod.	<1.0	
II. Sodium-Voided Configurations.	A. Sodium Voiding in Central 9 Drawers (r = 9.35 cm).	Slightly <1.0	
	a. Core Loading Adjusted So That All Steps Subcritical Except As Noted.		
	1. Void Upper Axial Blanket.		
	2. Void Core and Blanket Above Midplane.		
	3. Void Full Height of Core and Blankets.		
	B. Sodium Voiding in Central 21 Drawers (r = 14.28 cm). Leave Central 9 Drawers Voided.	Slightly <1.0	
	1. Void Upper Axial Blanket.		
	2. Void Core and Blanket Above Midplane.		
	3. Void Full Height of Core and Blankets.		
	C. Sodium Voiding in Central 37 Drawers (r = 18.96 cm). Leave Central 21 Drawers Voided.	Slightly <1.0	
	1. Void Upper Axial Blanket.		
	2. Void Core and Blanket Above Midplane.		
	3. Void Full Height of Core and Blankets.		
	4. Adjust Case C(3) to Critical.	1.0	Yes
	5. Measure Subcriticality of System with Central B <sub>4</sub> C Control Rod.	<1.0	

TABLE VI. (Contd.)

Phase	Step	Approximate $k_{eff}$ of System	Measurements at Critical (See Table II)
III. Fuel-Slump-Out Configurations.	A. Fuel Slump Out in Central 9 Drawers. All Fuel in Region $Z = 0$ to $Z = H/2$ Slumps in Region for $H/2$ to $H$ , where $H$ is the Half Height of Core and $Z = 0$ is at the Core Midplane.		
	1. Slump Upper Half of Core.	<1.0	
	2. Slump Over Full Core Height.	<1.0	
	B. Fuel Slump Out in Central 21 Drawers. Leave Central 9 Drawers Slumped.		
	1. Slump Upper Half of Core.	<1.0	
	2. Slump Over Full Core Height.	<1.0	
	C. Fuel Slump Out in Central 37 Drawers. Leave Central 21 Drawers Slumped.		
	1. Slump Upper Half of Core.	<1.0	
	2. Slump Over Full Core Height.	<1.0	
	3. Adjust Case C(2) to Critical.	1.0	Yes
	4. Measure Subcriticality of System With Central $B_4C$ Control Rod.	<1.0	
	5. Measure Subcriticality of System With Layer of $B_4C$ on Top of Lower Fuel Slump Region.	<1.0	
	6. Measure Subcriticality of System With Layer of $B_4C$ on Top of Upper Fuel Slump Region.	<1.0	

TABLE VI. (Contd.)

Phase	Step	Approximate $k_{\text{eff}}$ of System	Measurements at Critical (See Table II)
	7. Start From Step C(2). Unslump Fuel in Bottom Half of Core.	Near 1.0	
	8. Measure Subcriticality of System with Central B <sub>4</sub> C Control Rod.	<1.0	
	D. Move Fuel Slump Regions into Axial Blankets.		
	1. Upper Half of Core (with Bottom Half Unslumped).	<1.0	
	2. Full Core Height.	<1.0	
	3. Adjust D(2) to critical.	1.0	Yes
	4. Measure Subcriticality of System with Central B <sub>4</sub> C Control Rod.	<1.0	
IV. Fuel-Slump-In Configurations.	A. Fuel Slump In in Central 9 Drawers. All Fuel in Region $Z = H$ to $Z = H/2$ Slumps into Region $Z = H/2$ to $Z = 0$ , where $H$ is the Half Height of the Core and $Z = 0$ is at the Core Midplane.		
a. All Measurements With Central 37 Drawers Na-Voided.	1. Slump Upper Half of Core.	<1.0	
	2. Slump over Full Core Height.	<1.0	
b. All Fuel-Slump-Out Steps Removed.	B. Fuel Slump In in Central 21 Drawers. Leave Central 9 Drawers Slumped.		
c. Core Loading Ad- justed So That All Steps Subcritical Except As Noted.	1. Slump in Upper Half of Core.	<1.0	
	2. Slump over Full Core Height.	<1.0	
	C. Fuel Slump In in Central 37 Drawers. Leave Central 21 Drawers Slumped.		
	1. Slump in Upper Half of Core.	<1.0	
	2. Slump over Full Core Height.	Near 1.0	

TABLE VI. (Contd.)

Phase	Step	Approximate $k_{eff}$ of System	Measurements at Critical (See Table II)
	3. Adjust case C(2) to Critical.	1.0	Yes
	4. Measure Subcriticality of System with Central B <sub>4</sub> C Control Rod.		
	5. Measure Subcriticality of System with Layer of B <sub>4</sub> C on Top of Fuel Slump Region.		
	6. Start From Step C(3). Unslump Bottom Half of Core.	<1.0	
	7. Adjust Step C(6) to Critical.	1.0	Yes
	8. Measure Subcriticality of System With Central B <sub>4</sub> C Control Rod.		
V. Blanket Collapse Configurations	A. Start From Step IV C(6). Fuel Slumped in Top Half of Core and Unslumped in Bottom Half of Core.		
a. All Measurements With Central 37 Drawers Na-Voided	1. Collapse Upper Axial Blanket Material So That it is Directly on Top of Slumped-In Core Material.	<1.0	
b. Core Loading Adjusted So That All Steps Sub- critical Except As Noted.	2. Slump In Fuel in Bottom Half of Core.	<1.0	
	3. Collapse Lower Axial Blanket onto Slumped Fuel Region.	Near 1.0	
	4. Adjust Step A(3) to Critical.	1.0	Yes
	5. Measure Subcriticality of System with Central B <sub>4</sub> C Control Rod.	<1.0	

TABLE VII. Outline of Measurements for Safety-Related Critical Experiments

Phase/Step (See Table I)	$k_{\text{eff}}$	Spectrum	$\beta_{\text{eff}}$	Reaction Rate Profiles	Material Worths	Doppler Coefficient
I A (Ref. Core)	✓	Core Center	✓	Axial and Radial	Fuel, Structure, Control, Coolant -- Axial and Radial Profiles	Core Center
II C (4) (Na-Void)	✓	Core Center	✓	Axial and Radial	Axial Profiles	Core Center
III C (3) (Fuel-Slump-Out, 1)	✓			Axial and Radial	Axial Profiles	Fuel Region
III D (3) (Fuel-Slump-Out, 2)	✓			Axial	Axial Profiles	
IV C (3) (Fuel-Slump-In, 1)	✓	Core Center	✓	Axial and Radial	Axial and Radial Profiles	Core Center
IV C (7) (Fuel-Slump-In, 2)	✓			Axial	Axial Profiles	
V A (4) (Blanket Collapse)	✓	Core Center		Axial and Radial	Axial Profiles	Core Center

The program, as presented in Table VI, is divided into the following five major phases.

- I. Reference Core.
- II. Sodium Voided Configurations.
- III. Fuel-Slump-Out Configurations.
- IV. Fuel-Slump-In Configurations.
- V. Blanket Collapse Configurations (with Fuel-Slump-In).

In each phase the core perturbations have been accomplished in several axial and radial steps. In this manner data is provided to test both the magnitude of the perturbations and the trends of a perturbation as a function of other parameters. A central control rod has been included in several of the configurations in order to test the ability of the analysis to predict the perturbed case with control present.

The program of cores and measurements outlined in Tables VI and VII is estimated to take from 12 to 15 months. Subsequent to the preparation of this program plan a new schedule was developed for ZPR-9 which considerably reduced the anticipated length of the safety-related critical experiments. The development of a revised program plan aimed at a program of approximately 6 months duration is now necessary.

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4. *Physics of Reactor Safety*, Quarterly Report, January-March 1976, Argonne National Laboratory Report, ANL-76-72, pp. 9-12.
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